WC–Ni$_3$Al–B composites prepared through Ni+Al elemental powder route

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Abstract: In order to develop the liquid phase sintering process of WC–Ni$_3$Al–B composites, the preparation process of WC+Ni$_3$Al prealloyed powder by reaction synthesis of carbonyl Ni, analytical purity Al and coarse WC powders was investigated. DSC and XRD were adopted to study the procedure of phase transformation for the 3Ni+Al and 70%WC+(3Ni+Al) mixed powders in temperature ranges of 550–1200 °C and 25–1400 °C, respectively. The results demonstrate that the formation mechanism of Ni$_3$Al depends on the reaction temperature. Besides WC phase, there exist Ni$_2$Al$_3$, NiAl and Ni$_3$Al intermetallics in the powder mixture after heat treatment at 200–660 °C, while only NiAl and Ni$_3$Al exist at 660–1100 °C. Homogeneous WC+Ni$_3$Al powder mixture can be obtained in the temperature range of 1100–1200 °C. The WC–30%(Ni$_3$Al–B) composites prepared from the mixed powders by conventional powder metallurgy technology show nearly full density and the shape of WC is round. WC–30%(Ni$_3$Al–B) composites exhibit higher hardness of 9.7 GPa, inferior bending strength of 1800 MPa and similar fracture toughness of 18 MPa·m$^{1/2}$ compared with commercial cemented carbides YGR45 (WC–30%(Co–Ni–Cr)).

Key words: WC–Ni$_3$Al–B composites; Ni$_3$Al intermetallic; liquid phase sintering; mechanical properties

1 Introduction

Cemented carbide, such as WC–Co, is generally considered an adequate combination of strength and hardness [1]. This is one of the important reasons for the selection of these materials for a large number of industrial applications, including cutting tools, wear parts, extrusion dies and mining equipment [2]. Metals, such as Fe, Ni and Co, have been used as ductile second phases to improve fracture toughness, but they have poor adaptation in aggressive environments [3], which limits the use of WC-based composites for high temperature critical applications. A lot of research about Ni$_3$Al taken as alternative binder for cemented carbide has been investigated [3–5]. This is mainly due to the properties [6,7] that Ni$_3$Al possess, such as increasing strength with temperature to a maximum at 700–800 °C, good oxidation resistance and good resistance to aqueous acidic corrosion environments. Meanwhile, with the addition of small amounts of boron, the ductility of Ni$_3$Al can be greatly improved [8]. At the same time, the wettability of Ni$_3$Al on WC and TiC is sufficient and the possibility of forming compacts of WC/TiC–Ni$_3$Al has been reported [9]. These properties make Ni$_3$Al a potentially attractive replacement for cobalt in the fabrication of both titanium and tungsten carbide matrix composites [4].

Liquid phase sintering [10], hot-pressing [4] and melt infiltration [11] are the three common production techniques for the processing of WC–Ni$_3$Al and TiC–Ni$_3$Al composites with promising mechanical properties comparable to that of commercially available WC–Co and TiC–Ni cerments, but the WC–Ni$_3$Al composites form a semi-continuous second phase with some remnant Ni$_3$Al-rich areas owing to the relatively large size of the starting Ni$_3$Al prealloyed powder, which may decrease the performances of the composites. Ni$_3$Al prealloyed powder becomes an obstacle during the processing of these cerments by traditional liquid phase sintering techniques. However, little information has been published on the preparation of Ni$_3$Al prealloyed powder by reaction synthesis from powder state specimens instead of compaction specimens [12]. Moreover, the effect of temperature on phase transition during the synthesis from powder state specimens is not

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very clear. Further studies on the processing of these composites by traditional liquid phase sintering are still necessary.

In this study, the preparation process of WC+Ni$_3$Al prealloyed powder by the addition of ceramic particles to the Ni+25%Al (molar fraction) powder mixture (abbreviated as 3Ni+Al in the text) was presented. Furthermore, the effect of temperature on phase transition in the 70%WC+(3Ni+Al) (mass fraction) mixed powders was investigated in detail, and the thermal effect of the 3Ni+Al powder mixture was studied. In addition, the mechanical properties of WC−30% (Ni$_3$Al−B) (volume fraction) composites prepared by liquid phase sintering through WC+Ni$_3$Al prealloyed powder were compared with commercial cemented carbides YGR45 under similar processing conditions.

2 Experimental

Commercially available pure carbonyl Ni powder (purity 99.80%), coarse WC powder (purity 99.50%) and analytical purity Al powder (purity 99.30%) with average particle sizes of 3, 10 and 15 µm, respectively, were used as starting materials. Figure 1 shows the SEM images of the powders. To obtain 3Ni+Al or 70%WC+(3Ni+Al) mixed powders, Ni and Al powders or Ni, Al and WC powders were ball-mixed in a stainless steel container for 4 h. Stainless steel balls with diameter of 20 mm were used. The mass ratio of ball-to-powder was 1:1.

Thermal analysis was performed on 3Ni+Al mixed powders on a differential scanning calorimeter (DSC, Netzsch STA 449C). The heating was conducted in argon at a constant rate of 10 °C/min from 25 to 1400 °C. The 70%WC+(3Ni+Al) mixed powders were heated in a vacuum furnace with a vacuum degree of 1 Pa at 550, 750, 900, 1000, 1100 and 1200 °C, respectively, and dwell for 2 h, followed by furnace cooling to room temperature. The heating rate was 2 °C/min during the whole process in order to avoid possible SHS procedure. During the heat treatment, all the mixed powder specimens were degassed at 400 °C for 2 h in hydrogen atmosphere. After heat treatment, the morphologies of these powder mixtures were observed on the surface and the cross-section on an optical microscope. All these powder mixtures were ground and sieved to obtain fine powders. Phase identification was done by XRD with Cu K$_\alpha$ radiation.

WC−30% (Ni$_3$Al−B) composites were prepared from WC+Ni$_3$Al prealloyed powder and balanced coarse WC, Ni and B powders by conventional powder metallurgy technology. After sintering, the density was determined by the Archimedes' method. Vickers hardness was determined from polished sample by indentation hardness tester under load of 98 N. Fracture toughness was measured by indentation method [13]. The 3-point bending tests (span length of 15 mm) of the as-sintered samples (5.25 mm×6.5 mm×20 mm) were also done at a cross-head speed of 2 mm/min. The phase compositions were characterized by means of X-ray diffraction. The samples were cut, ground and polished with diamond paste, and examined on an optical microscope and scanning electron microscope (Model JSM−6701F, JEOL, Japan) with an X-ray energy dispersive spectroscopy (EDS) system.

3 Results

3.1 Differential scanning calorimeter analysis

A simultaneous differential scanning calorimeter (DSC) was used to determine the procedure of the reactions. Since the starting exothermic or endothermic
temperature is not well defined, the temperatures corresponding to the peak values in the curve are reported for the reactions. It can be noted that 4 endothermic peaks at about 659, 959, 1035 and 1172 °C while 2 exothermic peaks at about 434 and 848 °C exist, as shown in Fig. 2.

![DSC curve of 3Ni+Al mixed powders](image)

**Fig. 2** DSC curve of 3Ni+Al mixed powders

### 3.2 X-ray diffraction analysis

For the X-ray diffraction analysis, a summary of the X-ray diffraction patterns obtained from the 70%WC+(3Ni+Al) mixed powders by reaction synthesis at different temperatures of 550, 750, 900, 1000, 1100 and 1200 °C is given in Fig. 3. As the temperature rises to 550 °C, peaks related to the formation of Ni$_2$Al$_3$, NiAl and Ni$_3$Al appear, but there are still Ni and Al peaks. According to the DSC analysis and the fact that the diffusion rate of Al is higher than that of Ni [14], aluminum-rich intermetallics, namely Ni$_2$Al$_3$, forms at the interface between Ni and Al powders by the solid state interdiffusion below the eutectic temperature of 640 °C. When the temperature reaches 750 °C, the Al and Ni$_2$Al$_3$ peaks are completely replaced by those corresponding to NiAl and Ni$_3$Al, and the Ni peaks weaken. As the temperature is raised to 900, 1000 and 1100 °C, the Ni and NiAl peaks become weak gradually. After being dwelt at 1200 °C, the Ni and NiAl peaks almost disappear and Ni$_3$Al becomes the major phase, showing that the reaction is completed and the specimens only consist of Ni$_3$Al and WC.

Figure 4 (a) represents the XRD pattern of the WC–30%(Ni$_3$Al–B) composites. The Ni$_3$Al peaks of WC–30%(Ni$_3$Al–B) composites become left shift compared with the 70%WC+Ni$_3$Al prealloyed powder (Fig. 4(b)) dwelt at 1200 °C for 2 h.

![X-ray diffraction patterns of WC–30%(Ni$_3$Al–B) composites (a) and 70%WC+Ni$_3$Al prealloyed powder (b)](image)

**Fig. 4** X-ray diffraction patterns of WC–30%(Ni$_3$Al–B) composites (a) and 70%WC+Ni$_3$Al prealloyed powder (b)

### 3.3 Morphology of WC+Ni$_3$Al mixture

Figure 5 illustrates the photos of 70%WC+ Ni$_3$Al powder mixture dwelt for 2 h at 1200 °C. The morphologies of the appearance and cross section are shown in Figs. 5(a) and (b), respectively, which exhibit uniformity.

![Photos of 70%WC+Ni$_3$Al powder mixture dwelt at 1200 °C for 2 h: (a) Appearance; (b) Cross-section](image)

**Fig. 5** Photos of 70%WC+Ni$_3$Al powder mixture dwelt at 1200 °C for 2 h: (a) Appearance; (b) Cross-section
Figure 6 shows the back-scattered images of the cross-sections of 70%WC+Ni3Al powder mixture dwelt for 2 h at 1200 °C. Different contrasts are seen from these back-scattered images, indicating different phases. EDS analysis was carried out to determine these phases and the results are demonstrated in Fig. 6(b). Glutinous Ni3Al slice appears on the large WC particles. The powder mixture is loose and large amount of pores (black regions in the micrograph) exist.

The WC+Ni3Al powder mixture can be obtained between 1100 and 1200 °C and it can be easily crumbled in a grinding mill. The surface topography of the WC+Ni3Al prealloyed powder after grinding is illustrated in Fig. 7. Homogeneous and fine (≤40 μm) WC+Ni3Al prealloyed powder can be observed.

3.4 Microstructure and mechanical properties

The backscattered electron micrographs of WC–30%(Ni3Al–B) and WC–30%(Co–Ni–Cr) samples are shown in Fig. 8. The WC (gray phase) has a relatively uniform distribution in the binder phase Ni3Al (dark area in Fig. 8(a)).

The mechanical properties of the sintered samples are listed in Table 1.

4 Discussion

4.1 Effect of temperature on phase transformation

According to the Ni–Al phase diagram in Fig. 9, the binary system exhibits five stable intermetallic compounds, namely, NiAl1, Ni2Al3, NiAl, Ni3Al3 and Ni2Al. Three of them are detected in this study. The reaction mechanism of the 70%WC+(3Ni+Al) mixed powders can be determined.

Table 1 Mechanical properties of WC–Ni3Al–B composite and WC–Co–Ni–Cr cemented carbide

<table>
<thead>
<tr>
<th>Alloy</th>
<th>Hardness (HV)</th>
<th>Bending strength/MPa</th>
<th>Fracture toughness/(MPa·m1/2)</th>
<th>Relative density after low-pressure sintering/%</th>
</tr>
</thead>
<tbody>
<tr>
<td>WC–30% (Co–Ni–Cr)</td>
<td>8.5</td>
<td>2500</td>
<td>20</td>
<td>≥99</td>
</tr>
<tr>
<td>WC–30% (Ni3Al–B)</td>
<td>9.7</td>
<td>1800</td>
<td>18</td>
<td>≥99</td>
</tr>
</tbody>
</table>
Fig. 9 Ni–Al phase diagram [15]

First, for 3Ni+Al mixed powders, two exothermic peaks are observed at about 434 and 848 °C, respectively. A comparison of the temperature of the exothermic peaks and the XRD analysis results at these temperatures suggests that the exothermic peak at about 434 °C mostly results from the formation of Ni$_2$Al$_3$ and the gradient of the curve increases before the appearance of the exothermic peak at about 434 °C (Fig. 2), which indicates that the exothermic reaction (solid diffusion) between solid Ni and Al powders has happened. However, the exothermic peak at about 848 °C may be due to a small amount of melt aluminum-rich (NiAl$_3$) phase and heat emission of further reaction. It is possible that there are so few aluminum-rich phases that can not be detected by instrument at 750 °C. Four endothermic peaks appear at about 659, 959, 1035 and 1172 °C. It is believed that 659 °C corresponds to the melting point of Al. There is no exothermic peak at around 640 °C in the DSC curve as mentioned in Ref. [16] due to the influences of particle size of Al powder and powder state specimens. The particle size of Al powder is bigger than that of Ni powder only when Al melts and begins to diffuse at a high rate, which promotes the reaction synthesis between Ni and Al in turn. The test specimens in the powder state instead of compaction ones can reduce the eutectic reaction degree. The endothermic peaks at about 959, 1035 and 1172 °C may be caused by further reaction in intermediate NiAl.

Second, for 70%WC+(3Ni+Al) mixed powders, WC acts as heat buffer and reduces the contact area between the elemental powders. Intermediates, such as Ni$_2$Al$_3$, NiAl and Ni$_3$Al, are generated during the reaction synthesis depending on the temperature. According to the thermal analysis curve (Fig. 2), Al and Ni peaks disappear at temperature higher than 660 and 1100 °C, respectively. As temperature increases, intermediates form at the interface between Ni and Al from 200 to 660 °C. A certain amount of intermediates are produced with less amount of Al remaining and interstitial space formed due to the different solid diffusion coefficients of Ni and Al. When the temperature is close to 660 °C, Ni$_3$Al and NiAl grow at the expense of Ni$_2$Al$_3$, Ni and Al based on the XRD analysis. Upon further heating, Ni$_3$Al and NiAl become the main phases when the temperature exceeds 660 °C. In addition, from 660 to 1100 °C, three-phase Ni, Ni$_3$Al and NiAl co-exists and Ni almost disappears at 1100 °C. When the temperature reaches 1200 °C, the specimen contains uniform Ni$_3$Al and WC, which indicates a complete reaction. Based on the above discussion, the formation sequence of the various phases of 70%WC+(3Ni+Al) mixed powders are described as follows:

\[
\begin{align*}
\text{Ni+Al} & \rightarrow \text{Ni}_2\text{Al}_3 \ (200–660 \ ^\circ\text{C}) \quad (1) \\
\text{Ni+Ni}_2\text{Al}_3 & \rightarrow \text{NiAl}+\text{Ni}_3\text{Al} \ (\text{about } \ 660 \ ^\circ\text{C}) \quad (2) \\
\text{Ni+NiAl} & \rightarrow \text{Ni}_3\text{Al} \ (660–1100 \ ^\circ\text{C}) \quad (3)
\end{align*}
\]

Finally, the main point here is that the results from the experiment agree with Ref. [16], which indicates the similar formation phenomenon.

4.2 Effect of liquid phase sintering on microstructure and properties

As it can be seen from Fig. 8, the shape of WC grain in WC–30%(Ni$_3$Al–B) composite becomes round, which is different from the facets shape of WC grain in WC–30%(Co–Ni–Cr). It may be related to the dissolution-precipitation in the sintering process. After liquid phase sintering, the XRD pattern of WC–30% (Ni$_3$Al–B) composites in Fig. 4 presents the Ni$_3$Al peaks are left shifted compared with 70%WC+Ni$_3$Al prealloyed powder, which may be caused by the atom solid-solution in binder Ni$_3$Al during sintering. As listed in Table 1, the density of the sample was measured above 99% of the theoretical density, which is consistent with the very little porosity observed in the microstructure. The sample of WC–30%(Ni$_3$Al–B) has a remarkably high microhardness of 9.7 GPa, inferior bending strength of 1800 MPa and similar fracture toughness of 18 MPa·m$^{1/2}$ in comparison to the commercial coarse grain cemented carbides YGR45 (WC–30%(Co–Ni–Cr), WC grain size of 2.4 μm).

5 Conclusions

1) The WC+Ni$_3$Al powder mixture can be successfully obtained in the temperature range of 1100–1200 °C and it can be easily crumbled into fine powder due to the formation of pores during the reaction synthesis and powder state specimens instead of compaction specimens.

2) The formation mechanism of Ni$_3$Al depends on the reaction temperature. Besides WC phase, there exist nickel aluminate phases of Ni$_2$Al$_3$, NiAl and Ni$_3$Al in the...
powder mixture after heat treatment at 200–660 °C, while only NiAl and Ni3Al exist at 660–1100 °C.

3) WC–30%(Ni3Al–B) composites are prepared from WC+Ni3Al prealloyed powder and balanced coarse WC, Ni and B powders by conventional powder metallurgy technology and the composites can achieve nearly full density. WC–30%(Ni3Al–B) composites exhibit higher hardness, inferior 3-point bending strength and similar fracture toughness in comparison to the commercial coarse grain cemented carbides YGR45 samples.

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References


Ni+Al 元素粉末法制备 WC–Ni3Al–B 复合材料

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摘 要: 为发展 WC–Ni3Al–B 复合材料的液相烧结制备技术, 研究由羰基金属间化合物; 液相烧结; 力学性能

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